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A layer obtained from an aqueous dispersion containing a silicon/titanium mixed oxide powder prepared by flame-hydrolysis

5 The invention provides a layer which contains silicon/titanium mixed oxide powder prepared by flame-hydrolysis and the preparation and use thereof.

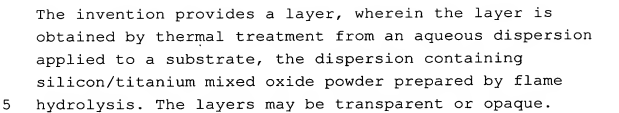
Layers which contain silicon dioxide and titanium dioxide are known. They are generally obtained by the so-called sol-gel process in which silicon and titanium alkoxides are deliberately hydrolyzed and polymerized in an organic solvent, usually an alcohol, and water. During polymerization a sol is initially produced and then, with increasing cross-linkage of the polymer units, a gel. This sol can be applied to a substrate, for example by dipcoating, and subsequently thermally treated and sintered.

It is also possible for the titanium component to be added in the form of titanium dioxide already and not as an alkoxide. Some of the silicon component may also be present as silicon dioxide.

Due to the high proportion of solvent in the gel, a high degree of shrinkage and the formation of cracks may also occur during extremely slow and careful drying. On the other hand it is difficult to form mechanically stable and thick layers with the low proportion of solids in the gel.

The object of the invention is to provide a layer on a substrate which has high mechanical stability and is free from cracks. The object of the invention is also a process for the preparation of this layer which leads to crack-free layers in a single coating step, wherein the thickness of the layer can be varied within wide limits.

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An aqueous dispersion is understood to be a dispersion with a proportion of powder of 0.1 to 60 wt.%, wherein the preferred range is between 2 and 40 wt.%. In addition, the dispersion may also contain inorganic or organic additives.

Thus, for example, the pH of the dispersion may be adjusted with tetramethylammonium hydroxide or hydrochloric acid. The dispersion can be prepared by methods known to a person skilled in the art such as, for example, by using an agitator, a rotor/stator, attrition mills or high-pressure homogenisers in which a predispersed stream under high pressure has a self-milling effect.

Flame hydrolysis is understood to be the hydrolysis of silicon and titanium compounds in the gas phase in a flame, produced by the reaction of hydrogen and oxygen. Here, highly disperse, non-porous primary particles are initially formed which can grow during further reaction to form aggregates and these can grow further to form agglomerates. Substantially spherical particles can also be obtained during synthesis, depending on the choice of reaction conditions.

A mixed oxide is understood to be an intimate blend of titanium dioxide and silicon dioxide at the atomic level, with the formation of Si-O-Ti bonds. In addition, the primary particles may also contain regions of silicon dioxide alongside titanium dioxide. To be differentiated from the mixed oxide powders used according to the invention are physical mixtures of silicon dioxide and titanium dioxide powders, the use of which is not suitable for preparing layers according to the invention. Also to be differentiated are sols and gels which may also contain

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Si-O-Ti bonds, but which are not suitable for forming layers according to the invention due to their porous structure and the low bulk density resulting from the method of preparation.

5 Furthermore, the powder may contain traces of impurities from the starting materials such as also impurities arising from the process itself. These impurities may amount to up to 0.5 wt.%, but generally not more than 100 ppm.

Silicon/titanium mixed oxide powders may be prepared, for example, by the method described in DE-A-4235996 by mixing silicon tetrachloride and titanium tetrachloride and burning these together with a hydrogen/air mixture. Furthermore, a silicon/titanium mixed oxide powder can be prepared by the method described in DE-A-19650500 with the proviso that the proportion of the mixed oxide component with the smaller proportion by weight, either silicon dioxide or titanium dioxide, does not exceed 20 wt.%. In the event that titanium dioxide is the component with the smaller proportion by weight, an aerosol obtained by nebulization and containing the solution or suspension of a salt of a titanium compound is fed into a gas mixture containing a silicon tetrahalide, hydrogen and air and is mixed homogeneously with this and then the aerosol/gas mixture is reacted in a flame inside a combustion chamber. In the event that silicon dioxide is the component with the smaller proportion by weight, the aerosol contains the salt of a silicon compound and the gas mixture contains a titanium tetrahalide.

Suitable substrates to which the layer may be applied, are borosilicate glass, silica glass, glass ceramics, materials with very low coefficients of expansion (ultra-low expansion, ULE, materials), or other inorganic substrates.

Thermal treatment is understood to be annealing in kilns, flame sintering or laser sintering.

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The layer may have a thickness between 100 nm and 1 mm, preferably between 1 μ m and 50 μ m and particularly preferably between 5 μ m and 15 μ m.

The BET surface area of the powder used for the layer may be between 5 and 500 m2/g and is controlled by adjusting the process parameters. Powders with BET surface areas between 20 and 50 m2/g are particularly advantageous.

The titanium dioxide content of the powder may be between 1 and 99 wt.%. A range between 2 and 12 wt.% is particularly preferred.

In accordance with a particular embodiment, the layer may contain a mixture of powders with high BET surface areas of at least 170 m2/g and those with low BET surface areas of at most 70 m2/g, preferably with high BET surface areas of at least 130 m2/g and those with low BET surface areas of at most 90 m2/g, wherein the ratio by weight of powders with lower to powders with higher BET surface areas is between 40:60 and 99.5:0.5.

The invention also provides a process for preparing the layer, wherein a dispersion which contains the silicon/titanium mixed oxide powder is applied to a substrate and this is then sintered by thermal treatment.

Borosilicate glass, silica glass, glass ceramics, materials with very low coefficients of expansion (ultra-low expansion, ULE, materials) or other inorganic substrates may be used as the substrate.

The dispersion may contain 0.1 to 60 wt.% of powder. A range between 2 and 40 wt.% is particularly preferred. A high density dispersion is preferred in particular for the application of thick layers. The dispersion may also contain inorganic or organic additives. Thus, for example, the pH of the dispersion can be adjusted with tetramethylammonium hydroxide or hydrochloric acid.

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The dispersion can be prepared by methods known to a person skilled in the art such as, for example, by means of an agitator, a rotor/stator, attrition mills or high-pressure homogenisers in which a predispersed stream under high pressure has a self-milling effect.

Application of the dispersion can be achieved by dipcoating, painting, spraying or spreading. Dip-coating is particularly preferred.

Thermal treatment may be performed, for example, by annealing in kilns, flame sintering or laser sintering.

Layers according to the invention can be used in composite materials with materials with very low coefficients of expansion (ultra-low expansion, ULE, materials), for photocatalytic applications, as coatings for self-cleaning mirrors (superhydrophilic constituents), for optical items such as lenses, as seals for gases and liquids and as mechanical protective layers.

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Examples

Example 1: Low surface area silicon/titanium mixed oxide powder

5.8 kg/h SiCl₄ and 0.37 kg/h TiCl₄ are evaporated together at about 170°C and fed into the central tube of a burner of 5 known construction. 3.7 Nm3/h of hydrogen and 6 Nm3/h of air and 0.65 Nm³/h of oxygen are also fed into this mixture. This gas mixture flows out of the burner nozzle and is burnt in the combustion chamber of a water-cooled flame tube. 0.5 Nm³/h of (secondary) hydrogen and 0.3 Nm³/h 10 of nitrogen are also fed to the jacket nozzle, which surrounds the central nozzle, in order to avoid caking. About 45 Nm³/h of air from the surroundings are also drawn into the flame tube which is under a pressure slightly lower than atmospheric. After flame hydrolysis, the 15 reaction gases and the pyrogenic silicon/titanium mixed oxide powder are drawn under suction through a cooling system by applying a reduced pressure and the particle/gas stream is thus cooled to about 100 to 160°C. The solids are separated from the exit gas stream in a filter or a 20 cyclone.

The pyrogenic silicon/titanium mixed oxide powder is produced as a white finely divided powder. Residues of hydrochloric acid which still adhere to the particles are removed from the mixed oxide powder at temperatures between 400 and 700°C, by treatment with steam-containing air, in a further step.

The BET surface area of the mixed oxide powder is $42 \text{ m}^2/\text{g}$. The analytical composition is $92.67 \text{ wt.} \% \text{ SiO}_2$ and $7.32 \text{ wt.} \% \text{ TiO}_2$.

The conditions of preparation are summarized in table 1 and further analytical data for the silicon/titanium mixed oxide powder obtained in this way are given in table 2.

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Example 2: High surface area silicon/titanium mixed oxide powder

5.8 kg/h SiCl₄ and 0.37 kg/h TiCl₄ are evaporated together at about 170°C and fed into the central tube of a burner of known construction. 2.0 Nm³/h of hydrogen and 7.7 Nm³/h of air are also fed into this mixture. This gas mixture flows out of the burner nozzle and is burnt in the combustion chamber of a water-cooled flame tube. 0.5 Nm³/h of (secondary) hydrogen and 0.3 Nm³/h of nitrogen are also fed to the jacket nozzle, which surrounds the central nozzle, in order to avoid caking. About 12 Nm³/h of air from the surroundings are also drawn into the flame tube which is under a pressure slightly lower than atmospheric. After flame hydrolysis, the reaction gases and the pyrogenic silicon/titanium mixed oxide powder are drawn under suction through a cooling system by applying a reduced pressure and the particle/gas stream is thus cooled to about 100 to 160°C. The solids are separated from the exit gas stream in a filter or a cyclone.

20 The pyrogenic silicon/titanium mixed oxide powder is produced as a white finely divided powder. Residues of hydrochloric acid which still adhere to the particles are removed from the mixed oxide powder at temperatures between 400 and 700°C, by treatment with steam-containing air, in a further step.

The BET surface area of the mixed oxide powder of Example 2 is $269 \text{ m}^2/\text{g}$. The analytical composition is $92.68 \text{ wt.} \% \text{ SiO}_2$ and $7.32 \text{ wt.} \% \text{ TiO}_2$.

The conditions of preparation are summarized in table 1 and further analytical data for the silicon/titanium mixed oxide powder obtained in this way are given in table 2.

Table 1: Experimental conditions for the preparation of pyrogenic silicon/titanium mixed oxide powder

Example		1	2
SiCl ₄	kg/h	5.8	5.8
TiCl ₄	kg/h	0.37	0.37
Primary air	Nm³/h	6	7.7
O ₂ added	Nm³/h	0.65	0
H ₂ core	Nm³/h	3.7	2.0
H ₂ jacket	Nm³/h	0.5	0.5
v _k (std)	m/s	23.9	22.6
gamma core	_	2.29	1.24
lambda core	_	1.03	1.61

Explanation: Primary air = amount of air in central tube (core); O_2 added = oxygen additionally fed into the core; H_2 core = hydrogen in the core nozzle; v_k (std) = speed of flow of gas leaving the core nozzle, corrected to standard conditions (273.15 K, 1 atm); gamma core = proportion of hydrogen in the core nozzle; lambda core = proportion of oxygen in the core nozzle. A detailed explanation of the expressions gamma and lambda and the calculation thereof is given in EP-A-0855368.

15 Table 2: Analytical data for the samples obtained in example 1 and example 2

Ex- ample	BET surface area m²/g	рН ⁽¹⁾	Bulk density g/l	Compacted bulk density	SiO ₂ content wt.%	TiO ₂ content wt.%
1	42	4.25	65	84	92.67	7.32
2	269	3.3	26	33	92.68	7.32

(1) 4% aqueous dispersion



Example 3: Preparation of an aqueous dispersion containing the mixed oxide powder from example 1

90 g of water and 10.0 g of mixed oxide powder from example 1 are dispersed in an agitator and treated for about 1 hour in an ultrasonic bath. Then the dispersion is filtered through a 60 µm sieve.

Example 4: Preparation of an aqueous dispersion containing the mixed oxide powders from examples 1 and 2

315 g of water and 10.5 g of mixed oxide powder from example 2 and 24.5 g of mixed oxide powder from example 1 are dispersed using an agitator and treated in an ultrasonic bath for about 1 hour. Then the dispersion was filtered through a $60~\mu m$ sieve.

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Example 5: Dip-coating of borosilicate glass using the dispersion from example 4

The dispersion from example 4 is applied to a borosilicate glass using dip-coating. The speed of immersion in the dispersion is 3 mm/s, the stationary time in the dispersion is 5 s, the speed of withdrawal is also 3 mm/s. Drying is performed in air at room temperature.

Sintering of samples preheated to about 300°C is performed using a laser (CO_2 laser Rofin Sinar RS700 SM).

The laser power is 590 W at a mirror frequency of 540 rpm. The sample is moved along under the laser beam with a calculated speed of about 2.32 mm/s and cooled slowly after sintering (circulating air oven, kept at 500°C for 1 hour and then cooled to room temperature over the course of 10 hours).

The thickness of the transparent layer was about 2.4 $\mu\text{m}\text{,}$ as determined by measuring the profile. The layer exhibited no cracks under an optical microscope.

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